FIELD OF THE INVENTION

This invention is related to utilizing molecular biology and recombinant DNA technology to isolate and/or identify sequences from different plant families.

BACKGROUND OF THE INVENTION

References describing codon usage include: Carels et al., J. Mol. Evol., Vol. 46, pp. 45-53 (1998) and Fennoy et al., Nucl. Acids Res., Vol. 21, No. 23, pp. 5294-5300 (1993).

10 AP2 like proteins and genes of Arabidopsis re described in copending U.S. Application Nos. 08/700,152; 08/879,827; 08/912,272; and 09/026,039.

SUMMARY OF THE INVENTION

The present invention relates to a method of isolating a target polynucleotide from a target plant species that encodes a polypeptide exhibiting a desired degree of sequence identity to a conserved region of a template polypeptide from a template plant species. The method comprises:

- 20 (a) identifying the amino acid sequence of the conserved region in the template polypeptide;
 - (b) generating an oligonucleotide comprising a sequence wherein the sequence or its reverse complement encodes at least four amino acids of the conserved region identified in step (a), wherein.
 - (i) the nucleotide of the first and second position of at least three codons are the same as the

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corresponding nucleotides in the template polynucleotide encoding the template polypeptide; and

(ii) the nucleotide of the third position of the codon of step (i) is the same as the nucleotide at the third position of the most preferred codon of the second plant class, family, genera, or species for that amino acid in the portion of the conserved region;

further wherein the oligonucleotide preferably does not comprise homopolymers of more than four nucleotides; and the oligonucleotide is not degenerate;

- (c) providing a composition comprising the target polynucleotide;
 - (d) contacting the oligonucleotide and the target polynucleotide under conditions that permit hybridization and formation of a duplex.

Identification of target polynucleotide can be accomplished by detection of the duplex of step (d). Further, both single stranded and double stranded target polynucleotides can be generated from the duplex of step (d).

DETAILED DESCRIPTION OF THE INVENTION Definitions

The usage of the term "plant family" herein refers to the common nomenclature used to classify organisms, for example Liliaceae and Orchidaceae are plant families.

General Method

The present invention relates to a method of isolating 20 and/or identifying genes in nucleic acids from a target plant species related to a gene or corresponding cDNA or other nucleic acids from a template plant species.

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Preferably, the target and template plant species are from different plant families.

In another embodiment of the invention, the method includes identifying and/or isolating from a target plant species a target polynucleotide that encodes a conserved region that exhibits at least 70% sequence a conserved region encoded by the template polynucleotide from another plant species.

The target and template polynucleotides can be either RNA or DNA or derivatives thereof. The oligonucleotides to be utilized can be RNA, DNA, or derivatives thereof, such as protein-nucleic acids, (PNAs). The target polynucleotide can be isolated from cDNA or genomic libraries or fixed on microarrays and need not be isolated directly from the second or target plant organism. Such plant sequences can be first subcloned into intermediary vectors or organisms.

The method utilizes sequences from a conserved region of the polypeptide encoded by the template polynucleotide. A "conserved region" is a primary sequence within polypeptide that correlates to an in vitro activity, in vivo activity, or a secondary structure. For example, the active site of a serine protease exhibits a particular tertiary structure that is responsible for the activity of protein. That same tertiary structure can be encoded by way 25 of different amino acid sequences, but certain portions of the sequence tend to be the same among the variants. The amino acid sequence identity of conserved regions from related proteins can be as low as approximately 35%. even polypeptides that exhibit about 35% sequence identity 30 can be useful to identify a conserved region. such conserved regions of related proteins typically, exhibit at least 50% sequence identity; even more typically

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at least about 60%; even more typically, at least 70% sequence identity, more typically at least 80%, even more typically about 90% sequence identity.

A. Identifying Conserved Regions

Conserved regions can be identified by locating a primary sequence within the template polypeptide that:

- (i) is a repeated sequence;
- (ii) forms some secondary structure, such as helices, beta sheets, etc.
- 10 (iii) establishes positively or negatively charged domains;
- (iv) represent a protein motif or domain. example, the Pfam web site describing the consensus sequence for a variety of protein motifs and domains. The sites on 15 World Wide Web in the UK http://www.sanger.ac.uk/Pfam/ and in the US at http://genome.wustl.edu/Pfam/. For a description of information included at the Pfam database, see Sonnhammer et al., Nucl Acids Res 26(1): 320-322 (January 1, 1998); and 20 Eddy SR, Durbin (1997)Sonnhammer EL, R Pfam: Comprehensive Database of Protein Families Based on Seed Alignments, Proteins 28:405-420; Bateman et al., Nucl. Acids Res. 27(1):260-262 (January 1, 1999); and Sonnhammer et al., Proteins 28(3):405-20 (July 1997).
- 25 From this database, consensus sequences of protein motifs and domains can be aligned with the template polypeptide sequence to determine the conserved region.

In addition, conserved regions can be determined by aligning sequences of the same or related genes in closely 30 related plant species. Closely related plants species

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preferably are from the same family. Alternativelly, plant species that are both monocots or both dicots are preferred.

Sequences from two different plant species are adequate. For example, sequences from Canola and 5 Arabidopsis can be used to identify the conserved region. Such related polypeptides from different plant species need not exhibit an extremely high sequence identity to aid in determining conserved regions.

Even polypeptides that exhibit about 35% sequence identity can be useful to identify a conserved region. More typically, such conserved regions of related proteins exhibit at least 50% sequence identity; even more typically at least about 60%; even more typically, at least 70% sequence identity, more typically at least 80%, even more typically about 90% sequence identity.

Typically, the conserved region of the target and template polypeptides or polynucleotides exhibit at least 70% sequence identity; more preferably, at least 80% sequence identity; even more preferably, at least 90% sequence identity; most preferably at least 92, 94, 96, 98, or 99% sequence identity. The sequence identity can be either at the amino acid or nucleotide level.

Sequence identity can be determined by optimal alignment of sequences to compare by the local homology algorithm of Smith and Waterman, Add. APL. Math. 2:482 (1981), by the homology alignment algorithm of Needleman and Wunsch, J. Mol. Biol. 48:443 (1970), by the search for similarity method of Pearson and Lipman, Proc. Natl. Acad. Sci. (USA) 85: 2444 (1988), by computerized implementations of these algorithms (GAP, BESTFIT, BLAST, PASTA, and TFASTA in the Wisconsin Genetics Software Package, Genetics Computer Group (GCG), 575 Science Dr., Madison, WI), or by inspection.



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Given that two sequences have been identified for comparison, GAP and BESTFIT are preferably employed to determine their optimal alignment. Typically, the default values of 5.00 for gap weight and 0.30 for gap weight length are used.

"Percentage of sequence identity" is determined by comparing two optimally aligned sequences over a comparison window, wherein the portion of the polynucleotide sequence in the comparison window may comprise additions or deletions (e.g., gaps or overhangs) as compared to the reference sequence (which does not comprise additions or deletions) for optimal alignment of the two sequences. The percentage is calculated by determining the number of positions at which the identical nucleic acid base or amino acid residue occurs in both sequences to yield the number of matched positions, dividing the number of matched positions by the total number of positions in the window of comparison and multiplying the result by 100 to yield the percentage of sequence identity.

Alternatively, the polynucleotides of а conserved 20 region of closely related species will hybridize under stringent conditions wherein one of the polynucleotides is a probe to determine the conserved region. "Stringency" is a function of probe length, probe composition (G + C content), and hybridization or wash conditions of salt concentration, 25 organic solvent concentration, and temperature. is typically compared by the parameter " T_m ", which is the temperature which 50% at of the complementary The relationship of hybridization conditions to T_m (in ${}^{\circ}C$) is expressed in the mathematical equation

 $T_{\rm m} = 81.5 - 16.6 (\log_{10}[Na^{+}]) + 0.41 (\%G+C) - (600/N)$ (1)

where N is the length of the probe. This equation works well for probes 14 to 70 nucleotides in length that are identical to the target sequence. For probes of 50 nucleotides to greater than 500 nucleotides, and conditions that include an organic solvent (formamide) an alternative formulation for T_m of DNA-DNA hybrids is useful.

 $T_m = 81.5 + 16.6 \log \{ [Na^{+1}/(1 + 0.7[Na^{+}]) \} + 0.41(\%G + C) - 500/L 0.63(\%formamide)$ (2)

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where L is the length of the probe in the hybrid. (P. Tijessen, "Hybridization with Nucleic Acid Probes" in Laboratory Techniques in Biochemistry and Molecular Biology, P.C. vand der Vliet, ed., c. 1993 by Elsevier, Amsterdam.) With respect to equation (2), T_m is affected by the nature of the hybrid; for DNA-RNA hybrids T_m is $10-15^{\circ}$ C higher than calculated, for RNA-RNA hybrids T_m is $20-25^{\circ}$ C higher. Most importantly for use of hybridization to identify DNA including genes corresponding to a template sequence, T_m decreases about 1 °C for each 1% decrease in homology when a long probe is used (Bonner et al., J. Mol. Biol. 81:123 (1973)).

20 Equation (2) is derived under assumptions equilibrium and therefore, hybridizations according to the invention are most preferably performed under conditions of probe excess and for sufficient time achieve equilibrium. The time required to reach equilibrium 25 be shortened by inclusion of a "hybridization accelerator" such as dextran sulfate or another high volume polymer in the hybridization buffer.

When the practitioner wishes to examine the result of membrane hybridizations under a variety of stringencies, an efficient way to do so is to perform the hybridization under a low stringency condition, then to wash the hybridization

B. Generating an Oligonucleotide

Once a conserved region is identified, an oligonucleotide can be generated to isolate and/or identify a target sequence. This oligonucleotide is usually not degenerate. Preferably, the oligonucleotide comprises a sequence wherein it or its reverse complement encodes a portion of the conserved region.

The portion is at least 3 amino acids in length, more typically, 4 amino acids in length; more typically at least 6 amino acids, even more typically at least 10 amino acids. Usually, the portion is at least than 40 amino acids; more usually, at least 30 amino acids; even more usually, usually at least 20 amino acids in length. A preferred range is from 3 to 18 amino acids in length.

The choice of which portion of the conserved region to use is based on convenience. Preferably, the portion of the conserved region is chosen to minimize the number of amino acids that are encoded by four or more codons. For example, the number of alanines, arginines, glycines, leucines, prolines, serines, threonines, and valines is minimized.

The sequence of the oligonucleotide is designed using 25 the following criteria:

- (1) Amino acid sequence of the conserved region of a template polypeptide;
- (2) Preferred codon usage in the class, family, genera, or species of target plant species; and
- 30 (3) Polynucleotide sequence of the template polypeptide.

This preferred codon can be the most preferred of the plant class from which the target plant species belongs. For example, if the target plant species belongs to the dicot class, the preferred codon can be the one that is preferred by all dicots. Alternatively, the preferred codon can be one preferred in the family, genera, or species that the target plant species belongs. (The terms class, family, genera, and species is used in accordance with the accepted classification system of all organisms.)

One example is illustrated below:

Conserved Region (AA): ...Aaa₁ - Aaa₂ - Aaa_{3...}

Template Polynucleotide

encoding conserved region: $(N_1N_2N_3) - (N_4N_5N_6) - (N_7N_8N_9)$

Preferred Codons for

20 conserved regions in

target plant species: $(X_1X_2X_3)$ $(X_4X_5X_6)$ $(X_7X_8X_9)$

Oligonucleotide: $(N_1N_2X_3) - (N_4N_5X_6) - (N_7N_8X_9) ...$

The third position of the second most preferred codon is utilized if the first two positions of the template polynucleotide do not match the most preferred codon, but the template polynucleotide matches the first two positions of the second most preferred codon.

Further, the oligonucleotide sequence is chosen to avoid homopolymers of more than four nucleotides.

Preferably, a portion of the conserved region is chosen to

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prevent such homopolymers from occurring in the oligonucleotide. Homopolymers can be included in the oligonucleotide if such a stretch is found in the template sequence and is preferred by the target plant species codon usage.

A higher percentage of guanosines and cytosines are preferred in the oligonucleotide sequence when a monocot target polynucleotide is to be isolated or identified using a template polynucleotide from a dicot plant species. Thus, for example, a guanosine or cytosine is preferred at the third position of the codons in the oligonucleotide when isolating and/or identifying a target sequence from a monocot using an Arabidopsis sequence as a template polynucleotide.

In contrast, higher percentage of adenines and thymidines are preferred in the oligonucleotide sequence when a dicot target polynucleotide is to be isolated or identified using a template polynucleotide from a monocot plant species. Thus, for example, an adenosine or thymidine may be preferred at the third position of the codons in the oligonucleotide when isolating and/or identifying a target sequence from a dicot, such as Arabidopsis, using a monocot sequences from corn as a template polynucleotide.

Oligonucleotides of the invention are at least 12, 16, 18, 20, 25 30, 35, 40, 45 or even at least 50 nucleotides in length.

The sequence and length are chosen to generate an oligonucleotide that is capable of forming a detectable duplex with target nucleotides. The oligonucleotide can include additional nucleotides, for example inosine, that bind to sequences in the template that flank the portion of the polynucleotide encoding the conserved region to

stabilize the formed duplex. Additional non-plant polynucleotide sequences may be helpful as a label to detect the formed duplex as a primary site for PCR or to insert a restriction site for later cloning of the isolated plant sequences.

More than one oligonucleotide can be generated from the conserved region to be used in the identification and isolation procedures.

C. <u>Isolating and/or Identifying Target Polynucleotide</u> Sequences

The target polynucleotide sequence is isolated by contacting the oligonucleotide of the invention with a composition that comprises the target polynucleotide under conditions that permit hybridization and formation of a duplex. The duplex is then detected and the target polynucleotide can be isolated.

Exemplary procedures for identifying and/or isolating target polynucleotides that can be used include polymerase chain reaction (PCR), Southern hybridization, and polynucleotide capture.

Isolation and/or identification of a target polynucleotide can be performed using any number of oligonucleotides constructed using the instant invention.

For example, a single probe can be used in colony 25 hybridization assays to identify from of library of clones the particular clone or clones that contain the desired target sequence. Such techniques are known, for example, for bacterial, yeast, and viral clones. Further, a single probe can also be used to generate the target starting material 30 polynucleotides from a comprising plurality of polynucleotides, for example in nick

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translation or cDNA synthesis or random priming or end labeling.

Single probes can be used in gel isolation techniques, such as Southern or Northern hybridization for identifying polynucleotides that correspond to the target polynucleotide to be isolated. For example, inserts of a cDNA library comprising the target polynucleotide are separated by length and are bound to a solid support so as to preserve the separation. Next, the oligonucleotide can be labeled and used to identify the fragments that hybridize to the oligonucleotide. Hybridization and wash stringency can be varied as defined above, but preferably stringent conditions are used.

Alternatively, a single oligonucleotide can be bound to 15 solid support to isolate the desired polynucleotide. The solid support can be exposed to a plurality of polynucleotides. The solid support can capture those polynucleotides that hybridize to the oligonucleotide, and the unwanted polynucleotides can be washed away. 20 target polynucleotide can be released from the solid support and further characterized or inserted into a vector.

Other methods for capturing target polynucleotides to a solid support using an oligonucleotide are described in Li et al., U.S. Pat. No. 5,500,356; and Laffler et al., U.S. Pat. No. 5,858,652.

Oligonucleotides of the invention can be used as primers in PCR to amplify the desired target polynucleotide sequences from a plurality of polynucleotides, such as a sample of mRNA from a tissue or a cDNA library. The reaction is run using the oligonucleotides as primers and mRNA (or cDNA) or genomic DNA from the target species as a substrate. The PCR product can be inserted directly into a

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vector for further processing. Alternatively, gel electrophoresis or other separations can be performed on the PCR product and the target polynucleotide can be identified by Southern hybridization techniques for further characterization or final isolation.

Amplification methods using a single oligonucleotide based on the instant invention specific for the target polynucleotide can be used for isolation and/or identification. Such a technique is single-primer PCR (SPPCR). A description of the method is described in Screaton et al., Nucl. Acids Res. 21: 2263-2264 (1993).

Other methods of isolating target polynucleotides with a single gene specific primer are described in Frohman et al., Proc Natl Acad Sci U S A 85(23):8998-9002 (Dec. 1988) and Uematsu et al., Immunogenetics 34(3):174-8 (1991).

Also, non-specific primers comprising, for example, poly-A, poly-T, or cap sequences, can be used in conjunction with a specific oligonucleotide of the invention.

PCR amplification methods can be performed using either one or two specific oligonucleotides generated from the conserved region of the template polypeptide. Preferably, the primers generate a product that is longer than the total length of the primers. Typically, using two primers, the portions of the conserved regions that are encoded by the oligonucleotides or their reverse complements are separated by at least about 5 amino acids, more typically by at least about 30 amino acids, more typically by at least about 50 or 100 amino acids. In another acceptable arrangement, the oligonucleotides (or their reverse complements) represent a portion of two different conserved regions of a single polypeptide. Then the polynucleotide between the

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conserved regions, perhaps inclusive of one, or both of them, is amplified.

Nested primers can be used to PCR amplify the target polynucleotide sequences.

5 Compositions and methods for reverse transcriptasepolymerase chain reaction (RT-PCR) is another means of isolating and/or identifying target polynucleotides utilizing oligonucleotide primers of the invention. example Lee et al, WO9844161A1 by Applicant Life Technologies. 10

Other amplification techniques, such as rapid amplification of cDNA ends can be used to isolate full length genes. One such procedure is described in Fehr et al., Brain Res Brain Res Protoc 3(3):242-51 (Jan. 1999).

15 D. <u>Identifying Target Polynucleotides</u>

The oligonucleotides of the invention can be utilized to identify the sequence of the target polynucleotides. example, the oligonucleotides can be used in a modified PCR procedure to obtain the sequence of the target polynucleotide. See, for example, Mitchell et al., U.S. Pat. No. 5,817,797; Uhlen, U.S. Pat. No. 5,405,746; Ruano, U.S. Pat. No. 5,427,911; Leushner et al, U.S. Pat. No. 5,789,168; and

The isolated target polynucleotide can be used in any 25 sequencing procedure, such as the known dideoxy termination method and its modifications, to identify the specific sequences.

E. Further Isolation of Target Polynucleotides

When the sequence of the target polynucleotides is 30 identified, primers can be constructed using sequence from

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the very termini of the target polynucleotides to "primer walk" and obtain the remaining sequences of the gene of which the target polynucleotides are a portion. See, for example, Screaton et al., Nucl. Acids Res. 21: 2263-2264 (1993).

The target polynucleotide can also be used to identify clones or colonies in a library that comprise sequences from the same gene as the target polynucleotide.

PLANT FAMILIES

Any plant from the plant kingdom can be used as a source of target or template polynucleotides. Without limitation, any of the plants from the monocot class, Liliopsida or from the dicot class, Magnoliopsida are of interest. Any families from these classes that can be used in the instant invention, including without limitation:

Liliaceae, Orchidaceae, Poaceae, Iridaceae, Arecaceae, Bromeliaceae, Cyperaceae, Juncaceae, Musaceae, Ameryllidaceae, Ranunculaceae, Arecaceae; Musaceae; Brassicaceae; Rosaceae; Fabaceae; Magnoliaceae; Apiaceae; Solanaceae; Lamaiaceae; Asteraceae; Salicaceae; Cucurbitaceae; Malvaceae; and Graminaceae.

Of particular interest as plants species from the following genera, without limitation, Anacardium, Arachis, Asparagus, Atropa, Avena, Brassica, Citrus, Citrullus, Capsicum, Carthamus, Cocos, Coffea, Cucumis, Cucurbita, Daucus, Elaeis, Fragaria, Glycine, Gossypium, Helianthus, Heterocallis, Hordeum, Hyoscyamus, Lactuca, Linum, Lolium, Lupinus, Lycopersicon, Malus, Manihot, Majorana, Medicago, Nicotiana, Olea, Oryza, Panieum, Pannesetum, Persea, Phaseolus, Pistachia, Pisum, Pyrus, Prunus, Raphanus,

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Ricinus, Secale, Senecio, Sinapis, Solanum, Sorghum, Theobromus, Trigonella, Triticum, Vicia, Vitis, Vigna, and Zea.

EXAMPLES

The invention is illustrated by the following Examples. The invention is not limited by the Examples; the scope of the invention is defined only by the claims following.

Example 1: General Materials and Methods

PLANT DNAs

Plant DNAs were isolated according to Jofuku and Goldberg (1988); "Analysis of plant gene structure", pp. 37-66 in Plant Molecular Biology: A Practical Approach, C.H. Shaw, ed. (Oxford: IRL Press).

OLIGONUCLEOTIDES

Oligonucleotide primer pairs were selected from template Arabidopsis gene sequences using default parameters and the PrimerSelect 3.11 software program (Lasergene sequence analysis suite, DNASTAR, Inc., Madison, WI). Selected primer pairs were then used to generate PCR products utilizing genomic DNA from Brassica napus as a target plant species and polynucleotides. products were either sequenced directly or cloned into E. coli $\mathtt{TOPO}^{\mathtt{TM}}$ using the TAvector cloning system according to manufacturer's guidelines (Invitrogen, Carlsbad, CA). Nucleotide sequences of PCR products and/or cloned inserts were determined using an ABI PRISM@ 377 DNA Analyzer as specified by the manufacturer (PE Applied Biosystems, Foster City, CA) and compared to the template Arabidopsis gene sequence using default parameters and the SegMan 3.61 software program (Lasergene sequence analysis suite, DNASTAR, Inc., Madison, WI). Brassica napus gene regions

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of greater than or equal to 17 nucleotides in length and 70% sequence identity relative to the Arabidopsis gene were selected and the nucleotide sequences translated into the corresponding amino acid sequences using standard genetic codes. Using the deduced amino acid sequences, the corresponding sequences of triplet codons of the Arabidopsis gene region, class-, family-, and/or species-specific codon usage oligonucleotide primer pairs were designed for use in identifying similar gene regions that would encode identical peptides various unrelated plant genera. In all cases, the DNA sequence of a primer or its reverse complement would be identical to the sequence of triplet codons of the Arabidopsis gene sequence at nucleotide positions 1 and 2. In some cases the nucleotide at position 3 of a triplet codon would be identical to the Arabidopsis codon if that codon is preferentially used in a given plant genera and/or species as determined by published codon usage In other cases, position 3 would be selected (e.g., A, G, C, T) using genera- and/or species-specific codon usage tables such that the designated nucleotide together with nucleotides in positions 1 and 2 will form a triplet codon that will encode an amino acid that is identical to that encoded by the Arabidopsis triplet codon. In some of these cases, where there is an equal probability of using one codon or another that encodes the same amino acid but differs only at position 3, then the selection of an A, G, C, or \mathbf{T} residue will not generate a string homopolynucleotides more than four nucleotides.

PCR

A typical PCR reaction consisted of 1-5 µg of template plant DNA, 10 pmol of each primer of a selected primer pair, and 1.25 U of Taq DNA polymerase in standard 1X PCR reaction buffer as specified by the manufacturer (Promega, Madison, WI). PCR

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reaction conditions consisted of one (1) initial cycle of denaturation at 94°C for 7 min, thirty-five (35) cycles of denaturation at 94°C for 1 min., primer-template annealing at 58°C for 30 sec., synthesis at 68°C for 4 min., and one (1) cycle of prolonged synthesis at 68°C for 7 min.

A typical single primer PCR (SPPCR) reaction consists of 1-5 μ g of template plant DNA, 10 pmol of a selected primer, and 1.25 U of Tag DNA polymerase in standard 1X PCR reaction buffer as specified by the manufacturer (Promega, Madison, WI). PCR reaction conditions consisted of twenty (20) cycles of denaturation at 94°C for 30 sec., primer-template annealing at 55° C for 30 sec., synthesis at 72° C for 1 min., 30 sec., two cycles (2) of denaturation at 94°C for 30 sec., primer-template annealing at 30° C for 15 sec., 35° C for 15 sec., 40° C for 15 sec., 45° C for 15 sec., 50° C for 15 sec., 55° C for 15 sec., 60° C for 15 sec., 65° C for 15 sec., and synthesis at 72° C for 1 min., 30 sec., thirty (30) cycles of denaturation at $94^{\circ}\mathrm{C}$ for 30 sec., primertemplate annealing at 55° C for 30 sec., synthesis at 72° C for 1 min., 30 sec., followed by one (1) cycle of prolonged synthesis at 72°C for 7 min.

IDENTIFICATION OF RELATED GENE SEQUENCES

Selected primers and/or primer pairs were used in PCR or SPPCR reactions using genomic DNAs isolated from selected plant genera to generate PCR products. Alternatively, primers and/or primer pairs could be used in RT-PCR reactions using RNA isolated from selected plant genera to generate PCR products using standard published procedures. PCR products were analyzed by agarose gel electrophoresis according to standard procedures. Specific products were extracted from agarose gels and either sequenced directly using the selected primer(s) as sequencing primers or first cloned into E. coli using the TOPOTM TA vector cloning

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system according to manufacturer's guidelines (Invitrogen, Carlsbad, CA). Cloned inserts were sequenced using an ABI PRISM™ 377 DNA Analyzer as specified by the manufacturer (PE Applied Biosystems, Foster City, CA). The DNA sequences obtained were then analyzed using the MapDraw 3.15 software program (Lasergene sequence analysis suite, DNASTAR, Inc., Madison, WI). Both nucleotide and deduced amino acid sequences were then compared to the template Arabidopsis and Brassica napus gene and amino acid sequences using default parameters and the MegAlign 3.18 software program (Lasergene sequence analysis suite, DNASTAR, Inc., Madison, WI) to verify gene identity.

Alternatively, selected primers and/or PCR products could be used directly as gene probes to screen plant genomic or cDNA libraries for putative related genes in various genera and/or species. Cloned inserts identified in this way would be sequenced and the nucleotide and deduced amino acid sequences analyzed as described previously.

GENERATING PRIMER SEQUENCES USING METHOD AS EXAMPLE 2: DESCRIBED -- COMPUTER SIMULATION

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(A)

GENE:

AGAMOUS

FUNCTION:

TRANSCRIPTION FACTOR

DOMAIN:

MADS BOX

AA SEQUENCE:

Predicted NT:

GGG AGG GGC AAG AUC GAG AUC AAG CGC AUC GAG

Maize

GGG AGA GGC AAG AUC GAG AUC AAG CGC AUC GAG

32/33

28/33

10 Rice GGG AGG GGG AAG AUC GAG AUC AAG CGG AUC GAG

31/33

Arabidopsis

GGG AGA GGA AAG AUC GAA AUC AAA CGG AUC GAG

29/33

(B)

GENE:

APETALA1

15 FUNCTION: TRANSCRIPTION FACTOR

DOMAIN:

MADS BOX

AA SEQUENCE:

N R AGG AUC GAG AAC AAG AUC AAC AAG CAG GUG ACC UUC

Predicted NT:

Maize

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cgg Auc gag aac aag auc aac cgg cag gug acc uuc AGG AUC GAG AAC AAG AUC AAC CGG CAG GUG ACG UUC

Rice

34/36

(M) 29/36

33/36

Arabidopsis

AGG AUA GAG AAC AAG AUC AAA AGA CAA GUG ACA UUC

(R)30/36

(C)

GENE:

APETALA2

25 FUNCTION:

TRANSCRIPTION FACTOR

DOMAIN: AP2 DOMAIN

AA SEQUENCE: Predicted NT: \mathbf{E} S Н I

GGC AGG UGG GAG UCC CAC AUC UGG GAC UGC

Maize

GGC cGc UGG GAa UCC CAC AUC UGG GAC UGC

27/30

30 Arabidopsis

GGA AGA UGG GAA UCU CAU AUU UGG GAC UGU

23/30

Example 3: SPECIFICITY OF CODON ADJUSTED PRIMERS

The following example illustrates the specificity of codon adjusted primer pairs. Primers 1 and 2 represent primers directly of from taken the sequence the template polynucleotide. Primers 1' and 2' are primers wherein the sequence has been codon adjusted for monocots according to the invention. These primers were used to identify target polynucleotides from corn and pice.

Primer 1

10 AA SEQUENCE
Coding Sequence:
Primer 1 Sequence:

Primer 1' (Codon Adjusted Sequence):

Sequence Identity to Primer 1:

Primer 2

AA SEQUENCE Coding Sequence: Complement

20 Primer 2 Sequence:

Codon Adjusted Sequence: Complement

Primer 2' Sequence:

% sequence Identity to Primer 2:

D C G L Q V 5' G GAC TGT GGG AAA CAA GTT TA 3' 5' G GAC TGT GGG AAA CAA GTT TA 3'

5' G GAC TGC GGG AAG CAG GTG TA 3' 17/21

81%

K Y R G V T L 5' AAG TAT AGA GGT GTC ACT TTG CA 3' 3' TTC ATA TCT CCA CAG TGA AAC GT 5'

5' TG CAA AGT GAC ACC TCT ATA CTT 3'

5' AAG TAC AGG GGC GTC ACC TTG CA 3' 3' TTC ATG TCC CCG CAG TGG AAC GT 5'

5' TG CAA GGT GAC GCC CCT GTA CTT 3' 19/23

83%

PCR was performed as described in Example 1 using genomic DNA from Arabidopsis thaliana, Oryza sativa (rice) and Zea mays (corn) as a source for the desired target polynucleotides.

RESULTS AND CONCLUSIONS:

PCR-amplified products of the expected size were generated using primers 1 and 2 and Arabidopsis genomic DNA as a substrate. No products were obtained in reactions using either rice or corn genomic DNA substrate.

On the other hand, PCR-amplified products were generated using the codon adjusted primers 1' and 2' and corn genomic DNA as a substrate. No products were obtained in a reaction using Arabidopsis genomic DNA substrate. Together these results demonstrate the general utility of designing codon adjusted primers for use in isolating/identifying gene orthologs from different plant families.

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>OAT ADC PROTEIN 65 aa GGFDTAHSAARAYDRAAIKFRGLDADINFNLSDYEEDLKQVTNWTKEEFVHILRRQSTGFARGSS

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>RICE ADC PROTEIN 65 aa
GGFDTAHAAARAYDRAAIKFRGVEADINFNYSDYEEDMRQMKSLSKEEFVHVLRRQSTGFSRGSS

35 >WHEAT ADC PROTEIN 65 aa GGFDTAHAAARAYDRAA KFRGVDADINFNLSDYEDDMKQVKGLSKEEFVHVLRRQSAGFSRGSS

>MAIZE ADC PROTEIN 65 aa 45 GGFDTAHAAARAYDRAAIKFRGVDADINFNLSDYDDDMKQVKSLSKEEFVHALRRQSTGFSRGSS

D

EXAMPLE 5. USE OF SHORT CODON ADJUSTED PRIMERS

Oligonucleotides

Codon adjusted oligonucleotides were designed as described previously. Derivatives of oligonucleotide 2' were generated as shown above and used as primers in combination with oligonucleotide 1' in PCR reactions using plant genemic DNA from Zea mays (corn), Avena sativa (oat), and Triticum aestivum (wheat) as a source of target polynucleotides.

PCR

A typical PCR reaction consisted of 1-5 μ g of target plant DNA, 10 pmol of primer 1' and 10 pmol of a derivitive of primer 2', and 1.25 U of Taq DNA polymerase/in standard 1X PCR reaction buffer as specified by the manufagturer (Promega, Madison, WI). PCR reaction conditions consisted of five cycles (5) denaturation at 94oC for 2 mi/nutes, 94oC for 30 sec., primertemplate annealing at 65oC for 15 sec., 60oC for 15 sec., 55oC for 15 sec., 50oC for 15/sec., 45oC for 15 sec., 40oC for 15sec., and synthesis at 600C for 1 min., 30 sec., and twenty (20) cycles of denaturation at 94oC for 30 sec., primer-template annealing at 55oC for 30 sec., synthesis at 72oC for 1 min., 30 sec., thirty (30) ϕ ycles of denaturation at 94oC for 30 sec., primer-template amnealing at 50oC for 30 sec., synthesis at 68oC for 1 min., followed by one (1) cycle of prolonged synthesis at 68oC for 7 min/.

25 Primer 1

AA SEQUENCE

Coding Sequence:

Primer Sequence:

Primer 1' (Codon Adjusted Sequence):

D C G L Q V

5' G GAC TGT GGG AAA CAA GTT TA 3'

5' G GAC TGT GGG AAA CAA GTT TA 3'

5' G GAC TGC GGG AAG CAG GTG TA 3'

W III

30 Primer 2

AA SEQUENCE Coding Sequence:

Complement

Primer 2 Sequence:

Codon Adjusted Sequence:

Complement

Primer 2' Sequence:

RISZU2'-1 (5 CODONS)

RISZU2'-2 (5 CODONS)

RISZU2'-3 (4 CODONS)

RISZU2'-4 (3 CODONS)

K Y R G V T

5' AAG TAT AGA GGT GTC ACT TTG CA 3'

3' TTC ATA TCT CCA CAG TGA AAC GT 5'

5' TG CAA AGT GAC ACC TCT ATA CTT 3'

5' AAG TAC AGG GGC GTC ACC TTG CA 3'

3' TTC ATZ TCC CCG CAG TGG AAC GT 5'

5' 7G CAA GGT GAC GCC CCT GTA CTT 3'

5' G CAA GGT GAC GCC CCT GT 3'

5' GGT GAC GCC CCT GTA CT 3'

5' GT GAC GCC CCT GTA CT 3'

31

5' GT GAC GCC CCT GT

RESULTS AND CONCLUSIONS:

As described in Methods, primer 2' derivitives vary in length from 15-18 bp that could encode a peptide of 4-5 amino acids in length. Figure 3 shows that PCR-amplified products were generated using primer 1' and primer 2' derivitives 1, 2, and 3 and all three genomic DNAs as a source of target polynucleotides.

These results demonstrate that the method as described can utilize conserved regions of greater than or equal to 4 amino acids in length for use in isolating/identifying gene orthologs from different plant families.

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